THEFT

Perfluorovinylhalophoaphines

or φ -alkeryiphosphinous acid. For example, tetra-ethyldiamide of perfluorovinylphosphinous acid bp by-90° C at 11 mm; nD 1 =470) was obtained in 53.6% yield in the reaction:

CF₂-CFMgI + CIP $N(C_2H_2)_2$ 2 \rightarrow CF = CFP $N(C_2H_5)_2$ 2 + MgCII Similarly, distinylamide of di-(trifluorovinyl)-phosphincus acid (bp 60° C at 25 mm; r_D^{20} 1.4098) was obtained in 37 5% yield on redistillation of fraction 49-53° C received in the reaction:

 $2CF_2 = CFM_5I + Cl_2PN(C_2H_5)_2 \longrightarrow (CF_2 = CF)_2PN(C_2H_5)_2 + 2MgClI$

The fractional distillation must not be carried to completion as the residue decomposes explosively. It was shown further that amides of the type $R'P(NR_2)_2$ are decomposed by dry HCl and form primary and secondary

Card 2/4

PerfluorovinyThalophosphines

77300 sov/63-4-6-34/37

chlorophosphines. Decomposition of perfluorovinglphosphinuous tetracthyldlamide with dry HCl gave per-Thorovinyldichlorophosphine (yield 66%; bp 81.5-820 C;

 $n_{\mathbf{D}}^{19}$ 1.4412):

 cF_2 - $cFP[N(c_2H_5)_2]_2 + 4HC1 \longrightarrow cF_2$ - $cFPC1_2 + 2(c_2H_5)_2$ NH·HC1

Similarly, the decomposition of di-(trifluoroviny1)phosphinous diethylamide gave di-(trifluorgyinyl)-chlorophosphine (yield 60%; bp 94-95° C; np 1.4095;

 $(CF_2-CF)_2PC1$). Also ethyldichlorophosphine $(C_2H_5PC1_2)$ was synthesized. The first two chlorophosphines in reaction with antimonous fluoride were transformed into the corresponding perfluorovinylfluorophosphines,

colorless liquids easily rlaring up in air. Perrluorovinyldichlorophosphine thus gave perfluorovinyldifluorophosphine (yield 64%; bp 2-30 C):

Card 3/4

Perfluorovinylhalophosphines

 $3CF_2 - CFPCl_2 + 2SbF_3 \longrightarrow 3CF_2 - CFPF_2 + 2SbCl_3$

Similarly, di-(trifluorovinyl)-chlorophosphine gave di-(trifluorovinyl)-fluorophosphine (CF₂-CF)₂PF (yield 50%; bp 63-65°C). There are 3 references, 1 U.K., 1 German, 1 Soviet. The U.K. reference is: F. Bennett, H. Emeleus, R. Haszeldine, J. Chem. Soc., 1953, p 1565.

SUBMITTED:

June 1, 1959

Card 4/4

5 (3)

Sterlin, R. N., Li-Wei-Kang, AUTHORS:

sov/62-59-8-37/42

Knunyants, I. L.

TITLE:

Perfluorodivinyl Mercury

PERIODICAL:

Izvestiya Akademii nauk SSSR. Otdeleniye khimicheskikh nauk,

1959, Nr 8, p 1506 (USSR)

ABSTRACT:

It is reported that perfluorodivinyl mercury (C4F6Hg) was obtained

from perfluorovinyl magnesium iodide and mercury chloride in an ether solution at -10-5° as a colorless liquid with a disagreeable odor (slightly soluble in water). C₄F₆Hg reacts rather

easily with iodine while perfluorovinyl iodide is formed. The physical properties of C4F6Hg differ considerably from those of

the perfluoroalkyl mercury derivatives. These have a high melting point and are easily soluble in water. In comparison to the compounds investigated they are considered to be halogene derivatives of Hg whereas the former are designated vinyl derivatives of mercury in which the pseudohalogenous character of

the perfluorovinyl radical is not prominent. There is

1 reference.

Card 1/

6.3600

77072 SOV/62-59-12-16/43

AUTHORS:

Sterlin, R. N., Bogachev, V. E., Yatsenko, R. D.,

Knunyants, I. L.

。 "我们还是我们们就是我们就是我们的人,我们就是我们的人,我们就是我们的人,我们就是我们的人,我们就是我们的人,我们就是我们的人,我们是我们的人,我们是我们就是

TITLE:

Reactions of Fluoroolefins. Communication 10. Concerning the Dependence of Chemical Properties of Fluoroelefins

on Polarity

PERIODICAL:

Izvestiya Akademii nauk SSSR. Otdelenie khimicheskikh

nauk, 1959, Nr 12, pp 2151-2155 (USSR)

ABSIRACT:

2-Diethylamino-1,2,2-trifluoro-1-bromoethane (I)

(bp 58° at 17 mm) was obtained by shaking perfluorovinyl bromide with diethylamine, at room temperature, for 2 hours. On hydrolysis of (I) with water, the diethylamide of fluorobromoacetic acid (bp 93° at 4 mm) was obtained in 85% yield. It was shown that the reaction rate of addition of diethylamine to perfluorovinyl halides increases with increasing polarity of the olefin molecule. Diethylamine reacts vigorously with diethylamide of

Diethylamine reacts vigorously with diethylamide of fluoroacetic acid; it almost fails to react with the diethylamide of chloroacetic acid, and reacts very slowly

Ca: d 1/2

Reactions of Fluoroplefins. Communication 10. 77072 Contending the Dependence of Chemical 507/62-59-12-16/43 Properties of Fluoroplefins on Polarity

with 2-diethylamino-1,2,2-triflucro-1-bromo- (or chloro) -ethane. There is 1 figure; 4 tables; and 5 references, 1 German, 1 U.K., 3 U.S. The 4 U.S. and U.K. references are: R. N. Haszeldine, J. Chem. Soc. 4259 (1952), A. Giacomo, R. Swith, J. Am. Chem. Soc. 77, 774 (1954); G. Rigby, H. Schroeder, U.S. pat 2409315 (1946); Hurwitz, W. Miller, Abstracts of Papers 114th Meeting, J. Am. Chem. Soc. 41 (1948).

SHEMISTED:

Marsh 31, 1968

Card 2/2

STERLIN, R.N.; PINKINA, L.N.; YATSENKO, R.D.; KNUNYANTS, I.L.

Perfluorovinyl derivatives of arsenic and antimony. Khim.nauka
i prom. 4 no.6:800-801 '59.

(Arsenic compounds)

(Antimony compounds)

STERLIN, R.N.; PINKINA, L.N.; KNUNYANTS, I.L.; NEZGOVOROV, L.F.

Exchange of radicals in the series of perfluoroalkenyl derivatives of magnesiu. Khim.nauka i prom. 4 no.6:809-810 '59.

(MIRA 13:8)

(Magnesium organic compounds)

(Hadicals (Chemistry))

STERLIN, R.N.; YATSENKO, R.D.; PINKINA, L.N.; KMUHYANTS, I.L.

Perfluorovinylhalophosphines. Khim.nauka i prom. 4 no.6:810-811
'59.

(Phosphine)

(Phosphine)

TO THE SECOND PROPERTY OF THE SECOND PROPERTY

STERLIN, R.N. [translator]; KNUNYANTS, I.L., akademik, red.;
VITKOVSKIY, D.P., red.; RABINOVICH, F.V., red.; ZASUL'SKAYA,
V.F., tekhn.red.

[Modern experimental methods in organic chemistry] Sovremennye metody eksperimenta v organicheskoi khimii. Pod red. I.L. Knuniantsa. Moskva, Gos.nauchno-tekhn.izd-vo khim.lit-ry, 1960. 627 p. (MIRA 14:1)

(Chemistry, Organic-Experiments)

5.3630

1287, 2209, 1266

s/062/60/000/011/006/016 B013/B078

AUTHORS:

Sterlin, R. N., Yatsenko, R. D., Pinkina, L. N.,

Knunyants, I. L.

TITLE:

Perfluoro Derivatives of Nonmetals

PERIODICAL:

Izvestiya Akademii nauk SSSR. Otdelcniye khimicheskikh

nauk, 1960, No. 11, pp. 1991 - 1997

TEXT: The preparation of perfluoro derivatives of phosphorus, arsenic, and antimony is described. From the reaction of perfluory megnesium iodide with AsCl₃, PCl₃, and SbCl₃ in ether solution only tertiary de-

rivatives were obtained: tri-(trifluorovinyl)arsine, tri-(trifluorovinyl)phosphine, and tri-(trifluorovinyl)stibine. Primary and secondary derivatives were not formed in this process. Perfluorovinyl dichloroarsine was obtained by splitting 10-alkyl-5,10-dihydrophenarsazine with dry HCl (Ref.4). A corresponding perfluorovinyl derivative was obtained in a quantitative yield as a result of the reaction of perfluorovinyl magnesium iodide with adamsite. Perfluorovinyl chloroarsine was

Card 1/3

Perfluoro Derivatives of Nonmetals

\$/062/60/000/011/006/016 B013/B078

isolated in a practically quantitative yield by the reaction of CF2 - CFAs(C6Ha)2NH with liquid HCl. By treating the tetraethyldiamide of phosphorous acid chloride and the tetraethyldiamide of ethyl phosphinic acid with dry, gaseous HCl in xylol solution, phosphorus trichloride, and ethyldichlorophosphine, respectively, were obtained. From the reaction of perfluorovinyl magnesium iodide with the tetraethyldiamide of phosphorous acid chloride, the tetraethyldiamide of perfluorovinyl phosphinic acid was obtained. This was converted into trifluorovinyl dichlorophosphine by reaction with dry HCl in ether solution. By treating the latter with antimony trifluoride, perfluorovinyl difluorophosphine was obtained. In a similar manner, the diethylamide of di-(trifluorovinyl) phosphinic acid was obtained from (C2H5)2NPCl2 and perfluorovinyl magnesium iodide. By decomposing it with dry HOl, di-(trifluorovinyl)chlorophosphine was synthesized. By treating the latter with antimony trifluoride, di-(trifluorovinyl)fluorophosphine was obtained. As opposed to the trifluoromethyl derivatives of arsenic and phosphorus, the prepared tri-(trifluorovinyl) arsine and tri-(trifluorovinyl)phosphine do not

Card 2/3

Perfluoro Derivatives of Nonmetals

s/062/60/000/011/006/016 B013/B078

separate trifluor ethylene when heated. Thus, the perfluor ovinyl radical in the said compounds does not show any properties of pseudohalogens. Ye. P. Shcherbina and L. F. Razgovorov assisted in this work. There are 8 references: 2 Soviet.

SUBMITTED:

June 4, 1959

Card 3/3

STERLIN, R.N.; LI VEY-GAN; KNUNYANTS, I.L.

Reactions of perfluorodivinylmercury. Zhur.VKHO 6 no.1;108-109
161. (Mira 14:3)

STERLIN, R.N.; DUBOV, S.S.; LI VEY-GAN; VAKHOUCHIK, L.P.; KNUNYANTS, I.L.

Certain regularities in the series of porfluorovinyl derivatives of the elements of groups IV and V of the periodic table.

Zhur.VKHO 6 no.1:110-111 '61.

(Vinyl compounds)

(Vinyl compounds)

STERLIN, R.N.; LI VEY_GAN [Li Wei-kang]; KNUNYANTS, I.L., akademik

Electronegativity of the perfluorovinyl radical. Dokl. AN SSSR
140 no.1:137-140 S.O "fi. (MIRA 14:9)

(Vinyl compounds)

STERLIN, R.N.; DUBOV, S.S.

Infrared spectra of perfluorovinyl derivatives of elements. Zhur.
VKHO 7 no.1:117-118 '62. (MIRA 15:3)

(Vinyl compounds—Spectra)

DUBOV, S. S.; CHELOBOV, F. N.; STERLIN, R. N.

Mass spectrometric study of some vinyl and perfluorovinyl compounds. Zhur. VKHO 7 no.5:585 162. (MIRA 15:10)

(Vinyl compounds—Spectra)

DUBOV, S.S.; TETEL*BAUM, B.I.; STERLIN, R.N.

Nuclear magnetic resonance of some perfluorovinyl derivatives.

Zhur. VKHO 7 no.6:691-692 '62. (MIRA 15:12)

(Vinyl compounds—Spectra)

KNUNYANTS, 1.L.; STERLIN, R.N.; TYULBNEVA, V.V.; PINKINA, L.N.

Pseudohalide properties of perfluoroalkenyl radicals in esters of perfluoroalkenylphosphinic acids. Izv. AN SSSR. Otd.khim.nauk no.6:1123-1127 Je '63. (MIRA 16:7)

1. Institut elementoorganicheskikh soyedineniy AN SSSR. (Phosphinic acid) (Radicals (Chemistry))

			1	
7	AUTHOR: Gololobov, Yu. G.; Dmit Yu. H.; Knunyants, I. L.; Sterl TITLE: A method for producing No. 169118 SOURCE: Byulleten' izobreteniy TOPIC TAGS: fluorine compound, organic compound ABSTRACT: This Author's Certif fluorovinylalkylphoaphinates. with perfluorovinylmarcury dur- modification of this method in	in R. N. alkyltrifluorovinylalkylp i tovarnykh znakov, no. phosphonic acid, organo ficate introduces a metho	thosphinates. Class 12, 6, 1965, 26 metallic compound, mercury d for producing alkyltri- phonic acids are interacted	
	ASSOCIATION: none SUBHITTED: 20Feb64 NO REF SOV: 000 Cord - 1/1	ENCL: 00 OTHER: 000	SUB CODE: GC, OC	1

KNUMYANTS, I.L., TYULENEVA, V.V.; PERVOVA, Ye.Ya.; STERLIN, R.N.

Pseudophosphonium compounds from triethyl phosphite and perfluore-clefins. Izv. AN SSSR. Ser. khim. no.10:1797-1801 0 64. (MIRA 17:12)

1. Institut elsmentoorganicheskikh soyedineniy AN SSSR.

VERVETKINA, A.K., inzh.; KOLCHINSKIY, Yu.L., inzh.; NIKOLAYEVSKIY, Ye.Ye., inzh.; RODIONOVA, R.G., inzh.; RYAFOLOV, A.F., inzh.; SOKOL, I.A., inzh.; STERLHI, S.L., inzh.; EVDEL'NANT, L.B., inzh.; ORLOV, V.M., kand. tekhn. nauk, retsenzent; YURGEL', B.I., inzh., retsenzent; FOKE, V.Ya., inzh., nauchn. red.; VOLNYANSKIY, A.K., glav. red.; SUDAKOV, G.G., zam. glav. red.; IOSELOVSKIY, I.V., red.; MARKOV, I.I., red.; MEL'NIK, V.I., red.; ONKIN, A.K., red.; STAROVEROV,. I.G., red.; TUSHIYAKOV, M.D., red.; CHERLOV, A.V., red.

[Engineering pipelines for industrial enterprises] Tekhnologicheskie truboprovody promyshlennykh predpriiatii. Moskva, Stroiizdat, 1964. 2 v. (MIRA 17:12)

VERVEYKINA, A.K., inzh.; KOLCHINSKIY, Yu.L., inzh.; NIKOLAYEVSKIY, Ye.Ya., inzh.; RODIONOVA, R.G., inzh.; RYAPOLOV, A.F., inzh.; SOKOL, I.A., inzh.; STERLIN, S.L., inzh.; EYDEL'NANT, L.B., inzh.; ORLOV, V.M., kand. tekhn. nauk retsenzent: YURGEL', B.I., inzh., retsenzent; FOKIN, V.Ya., inzh.; inzh., retsenzent; FOKIN, V.Ya., inzh.; inzh., red.; VOLNYANSKIY, A.K. red.; MARKOV, I.I., red.; MEL'NIK, V.I., red.; ONKIN, A.K., red.; STAROVEROV, I.G., red.; TUSHNYAKOV, M.D., red.; CHERNOV, A.V., red.; SUDAKOV, G.G., red.; IOSELOVSKIY, I.V., red.

[Technological pipings in industrial enterprises] Tekhnologicheskie truboprovody promyshlennykh predpriiatii. Moskva, Stroiizdat. Pt.1. 1964. 784 p. (MIRA 18:9)

"大"。 经政治的现在分词有效的现在分词的现在分词

BOLTUKHIN, A.K.; STERLIN, S.Z.; MUSHTAYEV, A.F.; MOROZOV, I.I.; KUDINOV, V.A.; MONAKHOV, G.A.; AZAREVICH, G.M.; LAPIDUS, A.S.; PROKOPOVICH, A.Ye., redaktor; RZHAVINSKIY, V.V., redaktor izdatel stva; TIKHANOV, A.Ya., tekhnicheskiy redaktor

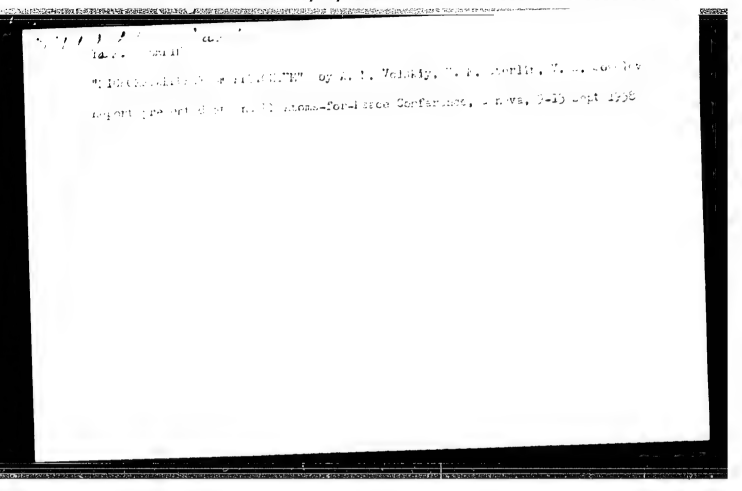
[Modernization of knee and column type milling machines; instructions]
Modernizataila konsol'no-frezernykh stankov; rukovodiashchie materialy.
Pod red. A.E.Prokopovicha. Moskva, Gos. nauchno-tekhn.izd-vo mashinostroit.lit-ry, 1957. 194 p.

(MLRA 10:8)

1. Moscow. Eksperimental'nyy nauchno-issledovatel'skiy institut metallorezhushchikh stankov (Milling machines)

KRASOV, Anatoliy Pavlovich; TROFIMOV, Arkadiy Alekseyevich; STERLIN, Ya.B., retsenzent; PESKOVA, L.N., red.; BOBROVA, Ye.N., tekhn. red.

[Journal-voucher accounting system on railroads] Zhurnal'noordernaia forma ucheta na zheleznykh dorogakh. Moskva, Vses.
izdatel'sko-poligr. ob"edinenie M-va putei soobshcheniia, 1961.
137 p. (MIRA 14:7)
(Railroads-Accounts, bookkeeping, etc.)



BOCHVAR, A.A., akademik, red.; YEMEL'YANOV, V.S., red.; ZVEREV, G.L., red. toma; IVANOV, A.N., red. toma; SOKURSKIY, Yu.N., red. toma; STER-LIN, Ya.M., red. toma; PEREVERZEV, V.V., red.; PCHELINTSEVA, G.M., red.; MAZEL', Ye.I., tekhn. red.

[Transactions of the International Conference On The Peaceful Uses of Atomic Energy] Trudy Vtoroy mezhdunarodnoy konferentsii po mirnomu ispol'zovaniyu atomnoy energii, 2d, Geneva, 1958. Izbrammye Doklady inos rannykh uchenykh. Moskva, Izd-vo Glav. uprav. po ispol'zovaniju atomnoi energ. pri Sovete Ministrov SSSR. Vol.6. [Nuclear fuel and ractor materials] IAdernoe goriuchee i reaktornye materialy. Pod obshchei red. A.A.Bochvara i Emel'ianova V.S. 1959. 702 p. (MIRA 14:10)

1. International Conference on The Peaceful Uses of Atomic Energy.
2d, Geneva, 1958. 2. Chlen-korrespondent AN SSSR (for Yemel'yanov).
(Nuclear fuels) (Nuclear reactors—Materials)

STERLIN, Yakov Moiseyevich, kand. tekhn. nauk; VOL'SKIY, A.N., akademik, red.; PANASENKOVA, Ye.I., red.; POPOVA, S.N., tekhn. red.

[Metallurgy of uranium] Metallurgiia urana. Pod obshchei red.
A.N.Vol'skogo. Moskva, Gosatomizdat, 1962. 418 p.

(MIRA 15:4)

(Uranium--Metallurgy)

- 1. SPERSIN, YE. A.
- 2. U JR (600)
- 4. Rells (Textile Machinery)
- 7. Methods for computing norms for reeling atuomats. Tekst.prom. 12 no. 11, 1952.

9. Monthly List for Mussian Accessions, Library of Congress, February 1953, Unclassified.

STERLIN, Ye.A., kundidat tekhnicheskikh nauk.

Meavers' werk organization. Tekst.prom.16 no.1:10-11 Ja '56.
(Meaving)

(MLRA 9:4)

Die entwicht: "Trye si atter af die Infloamme of Chomium tiend-lechnich instans di Inforfagen House in Ostron Crimina," owne w Textolo Trut, 30 May 4. 7: Tec em mo 1: 1mm, 'he', 1/17 (Project #17 36)

STERLIN. Yesim Abregovich; ZAMAKHOVSKIV, L.I., retsenzent; DOKHMAN, Ya.i., retheracent; SKGEL*, N.M., redsktor; DMITRIYEVA, N.I., tekhnicheskiy redsktor

[Principles of engineering standardization in textile manufacturing]
Osnovy takhnicheskogo normirovanila v tekstil*nom proizvodstva.

Moskva, Gos.nauchno-tekhn. izd-vo lit-ry po legkoi promyshl., 1957.

(M.Eh. 10:10)

177 p.

(Textile industry)

STERLIN, Ye.A., dots., kand.tekhn.nauk

Division of labor on automatic looms. Tekst. prom. 18 no.6;7-8

Je '58. (MIRA 11:7)

(Looms)

GOLUBEV. N., kand.tekhn.nauk; STERLIN, Ye., kand.tekhn.nauk; FEOKTISTOV,
M.; BREKHOV, A.; SIMAKIN, V.; KOZLOVA, L., tkachikha;
NIKONOVA, K.; CHERTKOV, L.; SLUTSKIN, S.; MINAYEV, I., insh.

Introducing a new organization of work; letter to the editor. Tekst.prom. 19 no.12:18 D '59. (MIRA 13:3)

1. Direktor Novo-Tkatskoy fabriki Glukhovskogo kombinata imeni V.I.Lenina (for Feoktistov). 2. Zaveduyushchiy 1-y tkatskoy fabrikoy kombinata "Vozhd' proletariata" (for Brekhov).
3. Nachal'nik tkatskogo proizvodstva fabriki im.M.V.Frunze (for Simakin). 4. Fabrika im. Frunze (for Kozlova, Nikonova).
5. Zaveduyushchiy normativno-issledovatel'skoy laboratoriyey po trudu fabriki im. M.V.Frunze (for Chertkov). 6. Zavedushchiy normativno-issledovatel'skoy laboratoriyey ramenskogo kombinata "Krasnoye Znamya" (for Slutskin).

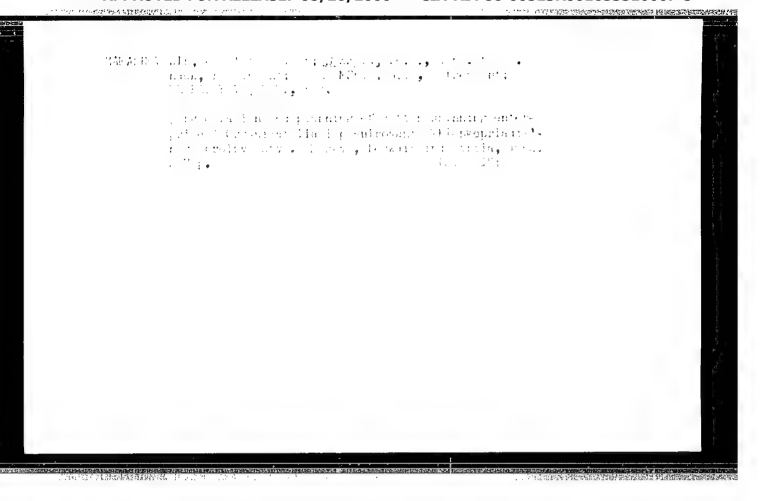
(Weaving)

STERLIN, Yefim Abramovich; POBEDIMSKIY, G.V., retsenzent; CHERTKOV, L.Ya., retsenzent; ZAMAKHOVSKIY, L.I., spets.red.; KOPELEVICH, Ye.I., red.; SHAPENKOVA, T.A., tekhm. red.

[Establishment of production norms in cotton spinning] Tekhnicheskoe normirovanie v khlopkopriadenii. Moskva, Izd-vo nauchno-tekhn.lit-ry RSFSR, 1961. 257 p. (MIRA 14:12) (Cotton manufacture—Production standards)

FIFELL, Yefim Abramovica; FEBEUTESKIY, G.V., retrengent; COFOLOVA, V.Yo., red.

[Work organization and the establishment of work norms for those who work at several machine tools in the textile industry] Organizatslia i normirovanie truda mnogostanochnikov v tekstil'noi promyshlennosti. Moskva, Izd-vo "leg-kain industriia," 1964. 197 p. (MIRA 19-7)



securing the scientific organization of the work of women spinning mechine operators. Tekst. prom. 25 no.10:1-4 0 '65. (MIRA 18:10) i. Khersonskiy filial Odesskogo tekhnologicheskogo instituta imeni lomonosova.

STERLIN, Ye.N.

Role of chronic bacterial carrier states in the pathogenesis of sporadic typhoid fever cases. Zhur. mikrobiol., epid. i immun. 40 no. 8:134-137 Ag '63. (MIRA 17:9)

1. Iz Meskovskogo instituta vaktsin i syveretek iseni Mechnikova.

POZHARSKIY, B.G.; STERLINGOVA, T.N.; PETROVA, A.Ye.

Hydrolysis and complex formation of uranyl in mineral acid solutions. Zhur. neorg. khim. 8 no.7:1594-1611 J1 '63.

(MIRA 16:7)

(Uranyl compounds) (Hydrolysis)

(Acids, Inorganic)

YAKOVLEV, Yu. V.; STERLINSKI, S.

Determination of contaminations in pure phosphorus by means of neutron activation, using X-ray spectrometry. Nukleon ka 7 no.3: 141-151 '62.

- 1. Institut geokhimii i analiticheskoy khimii AN SSSR, Dubna (for Yukovlev).
- 2. Institut yele nikh isstelestativ, TWI, Varshava, Otdeleniye analiticheskoy khimii (for Sterlinski).

24.6200

67357

21(1) AUTHORS: Sosnowski, R., Sterliński, S., Topa, J., Żylicz, J.

POL/45-18-6-3/5

TITLE:

Isomeric Transition in Hg 199

PERIODICAL:

Acta Physica Polonica, 1959, Vol 18, Nr 6, pp 573-580 (Poland)

ABSTRACT:

It was the aim of the present paper to investigate the spectrum of internal conversion electrons for the isomeric transition

in H_g^{199} from the $i_{13/2}$ to the $f_{5/2}$ -level. This 370-kev

transition was investigated under conditions, which permitted measurement of the ratio K/L and to estimate the E5 contribution. L.A.Sliv and A.M.Band had estimated the E5 admixture to 90%. Preparation of the Hg199 source is briefly described and shown in figure 1. For measurement of the internal conversion electron spectra, a magnetic spectrometer with a thick lens was used. A G-M counter of the BAT-10 type with a mica

window (1.3 $m_{\rm S}/{\rm cm}^2$) served as detector. The spectrometer had a resolution of 3.3%, the counting background did not exceed 3 counts/min; the electron absorption in the window was negligibly small. Measuring results are shown in several

Card 1/2

Isomeric Transition in Hg 199

67357 POL/45-18-5-3/5

diagrams. They show good agreement with those calculated theoretically for M4 transition in consideration of the finite nuclear dimensions and nuclear field shielding by the electron shell. The authors obtained: $K:L(M+N) = 1:(0.57\pm0.09)$: :(0.12 \pm 0.07). The mixture ratio of M4 to E5 is shown in figure 7. The maximum E5 admixture is found not to exceed 11%, which is in agreement with what was found by Pound and Wertheim. The authors finally thank Professor A. Soltan for his advice during construction of the spectrometer and for his keen interest. There are 7 figures and 13 references, 4 of which are Soviet.

ASSOCIATION:

Institute of Nuclear Research, Polish Academy of Science,

Varsaw

SUBMITTED:

April 25, 1959

Card 2/2

Gamma-spectroscopic ...

P/046/62/007/011/003/005 D256/D300

pulse-height analyzer. The two photoelectric peaks in the gamma spectrum corresponding to the 605 keV and 797 keV lines of Cs134 were used in the analysis; the identification of the lines was in addition verified by comparing the ratios of the areas under the peaks and measuring the rate of decay. The feasibility of the quantitative analysis was tested using samples of synthetic NaCl containing a known amount of CsCl, and the figure 2 x 10-3 ppm of Cs is quoted as the lowest amount which could be determined by this method under given conditions of activation and in the absence of any other long-lived impurity. It is pointed out that changes of the neutron flux in the reaction during the process of activation may produce an appreciable uncertainty of the results. There are 6 figures and 1 table.

ASSOCIATION:

Instytut Badań Jądrowych PAN, Warszawa, Zakład Chemii Analitycznej (Institute of Nuclear Research.

PAS Warsaw, Department of Analytical Chemistry)

SUBMITTED:

August. 1962

Card 2/2

L 13141-63

AFFTC/ASD BDS/EWT(m)

P/046/63/008/001/003/0014

AUTHOR:

TITLE:

Sterliński, Sławomir

Optimal irradiation and measurement conditions in an activation analysis. Gamma-spectrometric determination of trace contamination

in the presence of active matrix.

PERIODICAL:

Nukleonika, v. 8, no. 1, 1963, 57-67

The aim to reduce analytic activities in an activation analysis TEXT: leads to a gradual reduction of chemical operations and to optimum utilization of gamma-spectroscopic method. This method is limited by the fact that certain matrices are activated during the irradiation of the sample. Active matrices render the application of the gamma-spectroscopic method to direct sample analysis impossible. In this paper, gamma-spectroscopic determination of trace amounts of the element in presence of active matrix is considered. Formulae for relative statistical fluctuation of the number of counts due to the trace amount is offered. On the basis of these formulae optimal irradiation and measurement conditions are described. Several graphs illustrating the mathematical problem are included. Card 1/2/

Association: Nuclear Research Institute

STERLINSKI, Slawomir

Influence of the analyzer channel width on the shape of the photopeak of a monoenergetic gamma line. Nukleonika 8 no.10: 709-712 *63.

1. Department of Analytical Chemistry, Institut of Nuclear Research, Warsaw 9.

YURKENITE, Leopol's (Furkrewicz, Leopols), DTERLINGTH, Slovomir [Sterlinski, Slovomir]

Of thrum time of activity measurement in the presence of an unknown background. Pt. 1. Mukleonika 9 no.9:6972703 '64.

1. Institute of Muclear Research, Krakow Branch no.6 (for Jurkiewicz). 2. Institute of Muclear Research, Polish andemy of Sciences, Warsaw (for Sterlinski).

"APPROVED FOR RELEASE: 08/26/2000

CIA-RDP86-00513R001653310007-5

I. 21912-66 EVT(m)/EPF(n)-2
ACC FOR: 7.00111461

SOURCE CODE: PO/0046/65/010/011/0641/0659

AUTHOR: Sterlinski, Slawomir

28

ORG: Analytical Chemistry Department, Institute of Nuclear Research, Warsaw-Zeran

TITLE: Method for simultaneous counting of source and background for short-lived radioisotopes near the determination limit

SOURCE: Nukleonika, v. 10, no. 11, 1965, 641-659

TOPIC TAGS: scintillation spectrometer, radioisotope

ABSTRACT: The problem of simultaneously counting source and background radiations when determining trace amounts of short-lived radioisotopes by gamma scintillation spectrometry is discussed. In the case of time-consuming measurements (of the order of hours) the evaluation of the background on the basis of its counting in another time interval may prove to be false because of daily fluctuations. The error due to this may be in many cases eliminated by estimating background under a given photoelectric peak on the basis of the number of counts measured within another energy range of the spectrum. A method of estimating background within a measuring range is presented, the variance of the estimation is calculated, and a condition is pointed out under which the optimum (the minimum) relative accuracy is attained. The author thanks Mrs. Marecka and Mr. Kowalski for carrying out numerical calculations. Orig. art. has: 7 figures and 31 formulas. [NA]

SUB CODE: 18 / SUBM DATE: 21Jun65 / ORIG REF: 004 / OTH REF: 015

Card 1/1 1/1/25

SOURCE CODE: 70/00/0/00/011/000/0441/0453 Acc 1.7002.71 Company and the service of the servi C.C: Department of Analytical Chemistry, Institute of Nuclear Research, Warsaw-TYPE: Lower limit of detection for short-lived radioisotopes (Paper presented in the Conference on "Application of Thysico-Chemical Methods in Chemical Mallysis and in gualapest from 20 to 23 April 1966) SCURCA: Makkleonika, v. 11, no. 6, 1966, 441-453 TUPIC MAD: radioisotope, radioactivity measurement ADSTRACT: The relation of the "lower limit of detection" for short-lived radioisotopes and the measurement time and the so-called 'merit of the detector' was studied. It was found that the limit of detection reaches a minimum for the measurement time 1.81 $T_{\rm eff}$ (if the condition — 1, where $k_{\rm B}$ is the observed number counts of background, is fulfilled). The numerical value of the "lower limit" 1, where kn is the observed number of detection" can be calculated according to the included formulas. The theoretical conclusions are illustrated by two experiments using the radioisorones 66Cu (T₁ = 5.1 min) and ⁵²y (T₂ = 3.75 min). The author thanks Professor J. Minesewski for ellering the possibility of studies on this subject, Professor L. Juritaries (deconsect and Dr. R. Dybesynski for their discussions and remarks on this work, and rroledsor 3. Sicklerski for enabling the use of a neutron source. The author also thanks Miss M. Molbing, Miss T. Molesnik and Mr. J. Mowalski for their assistance in the measurements and calculations. 1/2 Card

L 19258-07
ACC NR: MF7002371

Grig. art. in Ing. Orig. art. has: 6 figures, 9 formulas and 2 tables. [Ni]
SUD CODE: 18 / SUDM DATE: 14Jan66 / ORIG REF: 003 / SOV REF: 001 / OTH REF: 014

ACC MR: AP7010679

SOURCE CODE: PO/0046/66/011/07-/0533/0553

AUTHOR: Scerlinski, Slavomiri; Dybezynski, Rajmund--Dybchinski R.

ONG: Department of Analytical Chemistry, Institute of Nuclear Research,

TITLE: Determination of cesium traces in mineral salts by means of neutron activation-ion exchange chromatography method

SOURCE: Nukleonika, v. 11, no. 7-8, 1966, 533-553

TOPIC TAGS: cesium, neutron activation analysis, ion exchange chromatography, isomer, chromatographic analysis, alkali metal, trace analysis

SUB CODE: 07,18

ADSTRACT: A neutron activation-chromatographic method for the determination of traces of cesium that makes use of the short-lived isomer 134mCs is presented. Cesium was isolated from macro amounts (0.5 g sample) of lighter alkali metals on the column with IR-3 cation exchanger (phenolsulfonic resin) before irradiation. After irradiation this cesium fraction was further purified from interfering activities by multistep ion exchange chromatography on micro-columns involving both cation and anion exchangers. The activity of the 31-keV photopeak was measured by gamma-spectrometric techniques using a very thin NaI(T1) crystal. The precision of the method (defined in terms of three standard deviations) was Cord 1/2

"APPROVED FOR RELEASE: 08/26/2000 CIA-RDI

CIA-RDP86-00513R001653310007-5

ACC NR: AP7010679

better than 12%. The sensitivity amounted to 4 x 10⁻¹¹ g Cs, i.e., 8 x 10^{-90%}. The method was shown to be fully applicable to the analysis of various mineral salts. The authors thank Mr. J. Kowalski for help in making the ion exchange separation, and Mr. R. Henrych for construction of the drop counter. Orig. art. has: 9 figures and 3 tables. [Orig. art. in Eng.] /NA

Card 2/2

STERLET, J. (a.

Sterlkov, S. Ya. - "The clinical treatment and therapy of Tirunea wounds in the constraint tesir possible.", Trudy Medinstituta (Izhev. gov. new. in-t), Vol. VI, 1907, . hh-h.

Co: U-hit, 17 Jety '3, (Letonia The nat 'nykn Statey, No. 19, 1969).

KAL'YU, r.I.; LUGINOVA, Ye.A.; IL'IN, S.Ye.; MATSKU, S.M.; ST.ML PAKH, O.N. (Moskva)

Structure and level of attendance of the rural population at medical and therapeutic institutions. Sov. zdrav. 20 no.7:17-22 '61. (MIAA 15:1)

1. Iz Instituta organizatsii zdravookhraneniya i istorii meditsiny imeni M.A.Semashko Ministerstva zdravookhraneniya SSSR.
(PUBLIC HEALTH, MUMAL)

STERIOGUY, C. L.

ilate, A. G., Sterioguy, C. L. and Paziellin, I. A.

(Moscow State Univ., and Frysical Inst., Acad. of Sci.)

J. Gen. Chem. (USSN) 14, 055-0 (1044) - English summary
Hydrocarbons of the cyclopentane series with a side-chain double bond.

/-cyclopentyle-2-butene and 3-cyclopentyl-1-butene.

Meat industry of Amur Province. Mias.ind. SSSR 2% no.6:39-40 '53. (MERA 6:12) 1. Upravlyayushchiy Amurskim myasotrestom. (Amur Province--Meat industry) (Meat industry--Amur Province)

Increasing the productive capacities. Miss.ind.SSSR 31 nr.l: 30 160. (MIRA 13:5) 1. Direktor Biyskogo myasokombinats. (Biysk--Packing houses--Equipment and supplies)

L 10658-63

EWP(q)/EWT(m)/BDS-AFFTC/ASD-JD/JXT(IJP)

ACCESSION NR: AP3001211

5/0078/63/008/006/1314/1319

AUTHOR: Mikheyeva, V. I.; Sterlyadkina, E. K.; Konstantinova, A. I.; Kryukova, O. N.

TITLE: Absorption of hydrogen by alloys of cerium with magnesium

SOURCE: Zhurnal neorganicheskoy khimii, v. 8, no. 6, 1963, 1314-1319

TOPIC TAGS: absorption; hydrogen; alloys; cerium; magnesium

ABSTRACT: Hydrogen absorption by cerium-magnesium alloys was investigated: alloys having more than 50 atm % Mg did not hydrogenate at room temperature; in alloys containing up to 30% Mg, the Ce appears to be preferentially hydrogenated; in the 30-50% Mg range, absorption corresponds to the formula CeH sub 3 • nMgH sub 2, in particular, MgH sub 2 • 2CeH sub 3 and Mg H sub 2 • CeH sub 3. Increase in Mg from 0-50% increases the induction period and hydrogenation time. If reaction is carried out at elevated temperature, alloys having up to 66% Mg can be hydrogenated but amount of H absorption is decreased. According to chemical and thermographic examination, the hydrogenation products are mechanical mixtures of the hydrides whose properties are distinct from those of MgH sub 2 and CeH sub 3 taken separately. Orig. art. has: 4 figures and 2 tables.

Card 1/2

"APPROVED FOR RELEASE: 08/26/2000

CIA-RDP86-00513R001653310007-5

L 10658-63

ACCESSION NR: AP3001211

2

ASSOCIATION: Institut obshchey i neorganicheskoy khimii im. N. S. Kurnakova, Akademii nauk SSSR (Institute of General and Inorganic Chemistry, Academy of Sciences, SSSR) Laboratoriya khimii gidradov i bora (Laboratory of Hydrides and Boron Chemistry)

SUBMITTED: 06Jul62

DATE ACQD: 01Jul63

ENCL: 00

SUB CODE: 00

NO REF SOV: 006

OTHER: 007

kes/2/2

L 13501-63 EWP(q)/EWT(m)/BDS AFFTC/ASD JD/JO

ACCESSION NR: AP3003482 S/0078/63/008/007/1710/1714

AUTHOR: Mikheyeva, V. I,; Sterlyadskhina, E. K.; Chertov, A. A.

TITLE: Hydrogenation of aluminum-cerium alloy

SOURCE: Zhumal neorganicheskoy khimii, v. 8, no. 7, 1963, 1710-1714

TOPIC TAGS: aluminum, cerium alloy, hydrogen, hydrogenation

ABSTRACT: Authors studied hydrogen absorption by cerium during its alloying with aluminum. Reason for this study was the marked absorption of hydrogen by cerium-magnesium alloy, characterized for a number of cases by whole and multiple proportions of CeH sub 3 to MgH sub 2. Metallic cerium, 99.9% pure aluminum and hydrogen which was obtained by the pyrolysis of titanium hydride, were used as materials in the study. The alloys were hydrogenated in accordance with the previously-described methodology (Mikheyeva and Kast, Zh, neorgan. khimii, 3, 1958, 260; Mikheyeva et al, Zh, neorg. khimii, 8, 1963, 1320) at room temperature and hydrogen pressure of about 1 atm. The hydrogen content in the hydrogenation products was determined by measuring the hydrogen volume during their reaction with diluted muriatic acid (1:5). The hydrogen volume which could be evolved during a reaction with metallic oxides, entering into the composition of the hydrogenation Card 1/2

 L 13501-63

ACCESSION NR: AP3003482

products, was calculated from the overall volume of evolved hydrogen. Authors found that alloying cerium with aluminum increases the induction period and hydrogenation time. The involvement of aluminum in the hydrogenation process was shown by observations over the hydrogenation process and analysis of the properties of the hydrogenation products. The maximum hydrogen absorption by aluminum corresponds to the empirical formula CeH sub 3 times 0.163AlH for an alloy with 14 atm. % of Al. Basically, the hydrogenation products of cerium-aluminum alloys evolve hydrogen in two stages, which, however, is not expressed as clearly as for CeH sub 3. Orig. art. has: 3 figures and 2 tables.

ASSOCIATION: Institut obshchey i neorganicheskoy khimii im. N. S. Kurnakova Akademii nauk SSSR (Institute of General and Inorganic Chemistry, Academy of Sciences, SSSR). Laboratoriya khimii gidridov i bora (Laboratory of hydride and boron chemistry).

SUBMITTED: 16Aug62

DATE ACQ: 02Aug63

ENCL: 00

SUB CODE: CH, ML

NO REF SOV: 004

OTHER: 008

Card 2/2

L 13500-63 EWP(q)/EWT(d)/EWT(m)/BDS AFFTC/ASD JD/JG ACCESSION NR: AP3003483 S,0078/63/008/007/1715/1721

AUTHOR: Mikheyeva, V. I.; Sterlyadkina, E. K.; Chertkov, A. A.

OD

TITLE: Hydrogenation of alloys of cerium with magnes tum, and aluminum 21

59

SOURCE: Zhurnal neorganicheskoy khimii, v. 8, no. 7, 1963, 1715-1721

TOPIC TAGS: cerium, magnesium, aluminum, hydrogenation, alloy

ABSTRACT: Authors studied the effects of aluminum on hydrogen absorption by cerium-magnesium alloys. The hydrogenation Zone of Ce-Mg-Al ternary alloys was determined at ordinary temperature and hydrogen pressure somewhat lower than atmospheric. The starting materials were metallic cerium, electrolytic magnesium and metallic aluminum of 99.9% purity. Dry hydrogen was obtained by decomposing titanium hydride which did not require any additional purification. The hydrogenation of the alloys was carried out on apparatus which did not basically differ from that described by Mikheyeva and Kost (Zh. neorgan. khimii, 3, 1958, 260). The hydrogenation zone of alloys in the system Ce-Mg-Al, rich in cerium, was determined at room temperature and normal pressure. It is distributed up to 25 atomic % of aluminum and 70 atomic % of magnesium. The maximum hydrogen absorption in the zone with 5-10% aluminum and 35-60% magnesium. It was shown that alloying melts in the Cord 1/2

。10.5000 Physics Christ Service Heart of Service 10.500 Physics Reference 15.000 Physics 15.000

L 13500-63 ACCESSION NR: AP3003483

Ce-Mg system with aluminum increases hydrogen absorption. During maximum hydrogenation of cerium to the composition CeH sub 3 and magnesium to the composition MgH sub 2, a part of the hydrogen is absorbed on account of being involved in the aluminum hydrogenation process. The presence of aluminum (5-15%) in alloys with a total content of aluminum and magnesium above 50% sharply reduces the hydrogenation induction period and produces a reaction which is less dependent upon the purity of the hydrogen. Aluminum also effects a lowering in the temperature for decomposing the cerium dihydride from 1080° (for pure dihydride) to 1010°. Orig. art. has: 5 figures and 1 table.

ASSOCIATION: Institut obshey i neorganicheskoy khimii im. N. S. Kurnakova (Institute of General and Inorganic Chemistry, Academy of Sciences, SSSB)

SUBMITTED: 14Aug62

DATE ACQ: 02Aug63

ENCL: 00

SUB CODE: CH, ML

NO REF SOV: 008

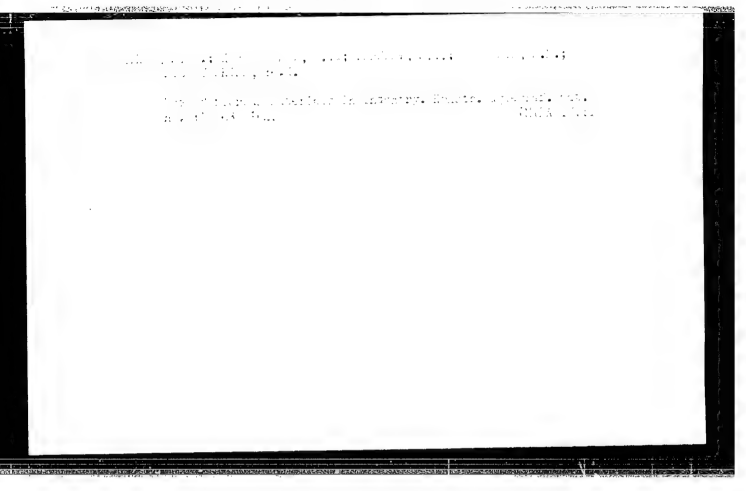
OFFIER: 007

Card 2/2

BULNEYATKINA, Ye. K.

STEPLYADKINA, Ye. K. "The Absorption by Glass of Radiations in the Infra-Red Portion of the Spectrum at High Temperatures." Min Construction Materials Industry USSR. All Union Sci Res Inst of Glass. Moscow, 1956. (Dissertation for the Degree of Candidate in Sciences)

So: Knizhnava Letopis', No. 17, 1956



STERLYADKINA, Z. K. (Aspirant)

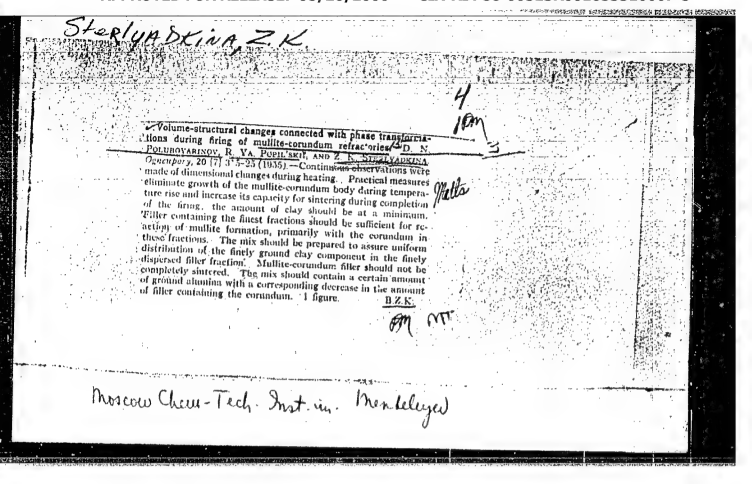
"An Investigation of the Volumetric-Structural Fluctuation Connected With Phase Transformations During the Firing of Mullite-Corundum Refractory Material." Cand Tech Sci, Moscow Order of Lenin Chemicotechnological Inst imeni D. I. Mendeleyev, 13 Dec 54. (VM, 2 Dec 54)

Survey of Scientific and Technical Dissertations Defended at USSR Higher Educational Institutions (12)

SO: SUM No. 556, 24 Jun 55

"APPROVED FOR RELEASE: 08/26/2000

CIA-RDP86-00513R001653310007-5



Wellyndkinn, Z.K

S/078/60/005/008/006/018 E004/B052

AUTHORS:

Mikheyeva, V. I., Sterlyadkina, Z. K., Kryukova, O. N.

TITLE:

Fusion Diagram of the System Aluminum - Copper - Lithium

PERIODICAL:

Zhurnal neorganicheskoy khimii, 1960, Vol. 5, No. 8,

pp. 1788-1795

TEXT: The authors first give a survey of the investigations published on the binary systems: Al - Cu (Ref. 3), Al - Li (investigated by F. I. Shamray and P. Ya. Sal'dau, Ref. 4), and Cu - Li (Refs. 5,6), and also a western paper (Ref. 7) on the ternary system. Then follows a description of their own method, the production of melts (analyses on Table 1), the thermal analyses by means of an N. S. Kurnakov pyrometer, and the investigation of the microstructure after etching with a 3 - 5% solution of HNO3 in alcohol (Fig. 4). The total fusion diagram of the system (up to 75 atom% of Cu+Li) is depicted in Fig. 5, while Fig. 1 gives a section of Al2Cu - AlLi. Fig. 2 of Al - Cu:Li = 4:1, and Fig. 3 of Al - Cu:Li = 9:1. Table 2 gives the data of the 15 points of the

Card 1/2

"APPROVED FOR RELEASE: 08/26/2000

CIA-RDP86-00513R001653310007-5

Fusion Diagram of the System Aluminum - Copper - Lithium

S/078/60/005/008/006/018 B004/B052

non-variant equilibrium. The system Al - Cu - Li is characterized by the crystallization of two ternary, incongruently melting phases: the S-phase consisting of the compound Al2CuLi, and the T-phase, whose composition in the homogeneity range approaches the compound Al6CuLiz. At 526°C, the ternary eutectic which corresponds to the common crystallization of the solid aluminum solution, also to the 0-phase (Al2Cu) and the S-phase (Al2CuLi), has the following composition: 73% of Al. 18.6% of Cu, and 8.4% of Li. All other non-variant points are transitional points. There are 5 figures, 2 tables, and 12 references: 6 Soviet, 2 US. 1 British. 2 German, and 1 Italian.

SUBMITTED:

May 27, 1959

Card 2/2

21332 S/078/61/006/004/001/018

B121/B216

11.2222

AUTHORS:

Dymova, T. N., Sterlyadkina, Z. K., Safronov, V. G.

TITLE:

A method for preparing magnesium hydride

PERIODICAL:

Zhurnal neorganicheskoy khimii, v. 6, no. 4, 1961, 763-767

TEXT: The present work discusses methods for the preparation of magnesium hydride and describes optimum conditions for a rapid and efficient preparation from the elements. Electrolytic magnesium of a purity of 99.3% and electrolytic hydrogen were used as initial materials. The synthesis was carried out in a rotating autoclave at 120-150 rpm, filled to one quarter with steel balls for grinding and mixing the material. The initial hydrogen pressure was 100-200 kg/cm². The resulting magnesium hydride was analyzed by measuring the hydrogen volume formed by reaction of magnesium hydride with a 5% solution of chromic anhydride. The reaction sets in at 260-270°C but comes to a stop when about 75% MgH2 has

formed because the magnesium becomes incrusted with the hydride. The yield was increased to 79% by applying a pressure of 200-300 kg/cm², increasing

Card 1/4

21332 S/078/61/006/004/001/018 B121/B216

A method for preparing ...

the temperature to 400-450°C and extending the reaction time to 15 hr. A yield of 96% magnesium hydride, leaving less than 1% unreacted magnesium was obtained by using 0.7% iodine as catalyst at a reaction temperature of 380-450°C and continuous grinding of the solid phases during 5-6 hr (Table). The reaction was also carried out by using carbon tetrachloride and a copper-magnesium alloy of the composition around the reagents at 420°C in the presence of Mg_Cu as activators. Grinding the reagents at 420°C in the presence of 1.5% CCl₄ yielded 85% magnesium hydride after 2 hr, and 100% magnesium hydride after 6 hr. The role of the activators is discussed. It is assumed that in the case of iodine catalyst an intermediate, magnesium subiodide, forms according to the reaction MgI₂ + Mg = 2MgI, which then reacts with hydrogen to form magnesium hydride. With carbon tetrachloride as activator, alkyl magnesium chloride is probably formed as well as magnesium subchloride. The authors thank V. I. Mikheyeva for discussion. There are 5 figures, 1 table, and 8 references: 3 Soviet-bloc and 5 non-Soviet-bloc.

Card 2/4

S/078/61/006/004/001/018 B121/B216

A method for preparing ...

Institut obshchey i neorganicheskoy khimii im. K. S. ASSOCIATION:

Kurnakova Akademii nauk SSSR (Institute of General and Inorganic Chemistry imeni N. S. Kurnakov, Academy of

Sciences USSR)

SUBLITTED:

November 1, 1960

Card 5/4

CIA-RDP86-00513R001653310007-5" APPROVED FOR RELEASE: 08/26/2000

21332 s/078/61/006/004/001/018 B121/B216

A method for preparing ...

Table: Temperature dependence of the magnesium hydride yields obtained by addition of iodine. Legend: 1) Mg in g; 2) reaction conditions; by addition in hr; 4) consumption of H2 in kg/cm²; 5) percentage in the product; 6) Mg_{met}; 7) calculated H₂ consumption

				Процент содержания в продукт		
Mg. #	(2) Bu	дерина диптивность.	Расход На ка/см ⁸	Mgli,	MEMEE (6)	MEO
100 100 100 100 100 100 200	200 250 300 350 380 420— -450 390—	- 0	25° 45 50 55 65 120°°	42,58 70,06 86,56 86,13 96,94 97,19	52,65 27,07 11,09 7,44 0,72 0,47 0,6	3.77 1,87 1,5 3.43 1,32 2,69

(7 Расчетный расход H₂: ° 60,3 кв/см²; °° 116,3 кв/см³.

card 4/4

CIA-RDP86-00513R001653310007-5" APPROVED FOR RELEASE: 08/26/2000

21333

S/078/61/006/004/002/018 B121/B216

11.2222

Dymova, T. N., Sterlyadkina, Z. K., Yeliseyeva, N. G.

AUTHORS: TITLE:

Some properties of magnesium hydride

PERIODICAL:

Zhurnal neorganicheskoy khimii, v. 6, no. 4, 1961, 768-773

TEXT: The present paper describes the behavior of magnesium hydride towards water and aqueous solutions. The magnesium hydride was prepared from the elements with addition of iodine (T. N. Dymova, Z. K. Sterlyadkina, V. G. Safronov, Zhurnal neorganicheskoy khimii, v. 6. no. 4, 1961, 763-767). [Abstracter's note: See abstract no. S/078/61/006/004/001/018]. This magnesium hydride was found to be comparatively unreactive. Only 56% of the hydride had hydrolyzed after 31 days (Fig. 2). Hydrolysis of magnesium hydride with solutions of sulfuric acid, chromic acid, potassium dichromate, ammonium chloride and cerium sulfate proceeds very rapidly. Treatment with aqueous solutions of sodium chlorate, boric acid, hydrogen peroxide, iodine in potassium iodide, and with slight amounts of cobalt and nickel lead to partial decomposition of magnesium hydride (Table 2). In a mixture of magnesium hydride, magnesium oxide, and metallic magnesium

Card 1/5

21333

s/078/61/006/004/002/018 B121/B216

Some properties of magnesium hydride

chromic acid selectively passivates the magnesium and quantitatively dissolves the magnesium hydride and magnesium oxide. A rapid method for determining magnesium hydride- magnesium oxide - magnesium metal mixtures was developed; it consists of measuring the volume of hydrogen liberated in sulfuric- and in chromic acid. The authors thank V. I. Mikheyeva for his valuable advice. There are 4 figures, 2 tables, and 10 references: 3 Soviet-bloc and 7 non-Soviet-bloc.

Institut obshchey i neorganicheskoy khimii im. N. S. ASSOCIATION:

Kurnakova Akademii nauk SSSR (Institute of General and

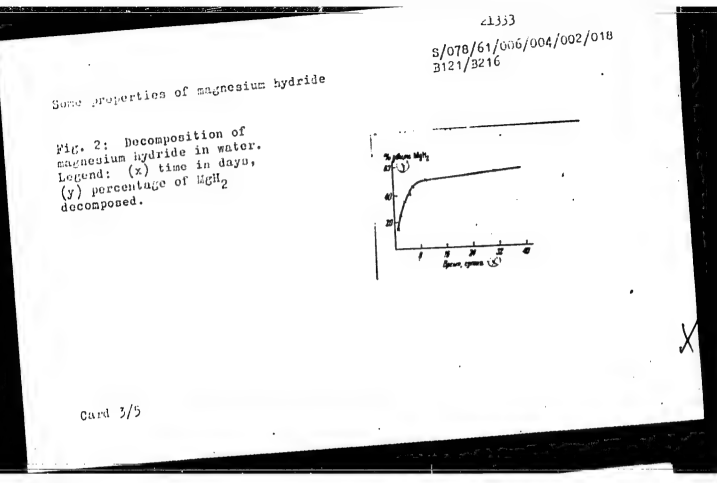
Inorganic Chemistry imeni N. S. Kurnakov, Academy of

Sciences USSR)

SUBMITTED:

November 1, 1960

Card 2/5



Some properties of magnesium hydride

Table ?: Influence of various substances on the hydrolysis of magnesium hydride and metallic magnesium. Legend: 1) substance; 2) concentration hydride and metallic magnesium; 5) magnesium hydride; in the solution; 3) quantity of hydrogen, ml, liberated by one gram of the substance in 15 min; 4) magnesium; 5) magnesium hydride; the substance in 15 min; 4) magnesium; 9) nickel salt; 6) average value; 7) no reaction; 8) cobalt salt; 9) nickel salt; 10) traces; 11) ditto; 12) theoretically, quantitative hydrolysis of 1 g of MgH₂ yields 1702.20 ml of H₂ and 1 g of Mg 921.60 ml of H₂.

Card 4/5

"APPROVED FOR RELEASE: 08/26/2000

CIA-RDP86-00513R001653310007-5

NALL OF	S/078/61/006/004/002/018 B121/B216	
Some propertie	es of magnesium hydriuc	
Table 2	Нещество Копцентра- ция раствора 1 2 среднее 1 1 2	
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_{Card} 5/5		

30029 5/020/61/141/001/013/021 B103/B147

5 2200

1043 1087 1273

Mikheyeva, V. I., and Sterlyadkina, Z. K.

AUTHORS:

Hydrogenation of cerium-magnesium alloys

TITLE:

Akademiya nauk SSSR. Doklady, v 141. no. 1, 1961, 104-106

FERIODICAL:

TEXT: Ce-Mg alloys were hydrogenated over a widem range of Mg concentrations than had been done by A. Sieverts and E. Roell (Zs. anorg. Chem., 146, 149 (1925)). Ce containing 0.75% of Nd, 0.4% of Pr, 1.0% of La, and 0.05% of Fe, and refined magnesium with a purity of at least 99.9% were used. The authors produced alloys with a Mg content of up to 85 at %, corresponding to "g Ce, in corundum crucibles under LiCl-KCl flux. They hydrogenated at room

temperature and a hydrogen pressure of 0.5 - 1 atm in an apparatus described by M. Ye. Kost, ZhNKh, 2, 2689 (1957). They used both cast and heat-treated samples (which absorb H2 much faster). H2 is absorbed by the alloys after

an induction period which increases considerably with increasing Mg content. but does not change in parallel with the composition. Samples nos. 13 - 15 (Table !) could only be hydrogenated after activation by previous heating. Alloys with more than 30% of Mg are split by hydrogenation along the cleavage Card 1/4

5/025/6:/141/001/013/021 B103/B147

Hydrogenation of cerium-magnesium...

Table ! shows the results. The initial range of the weak effect of Mg on H absorption of Ce is caused by the formation of solid solutions of Mg in Ce Maximum absorption is reached by alloys having a composition similar to MgCe. Peritectic reactions of formation of NgCe from Mg2Ce reduce H2 absorption more to alloys richer in Mg which, at room temperature, do not absorb hydrogen either in the initial state or after heat treatment. Up to 5 at% of Mg, Ce-Mg alloys hydrogenated up to saturation behave like CeH 3 From 10 at% of Mg conward, only vigorous pulverization causes inflammation whereas alloys with 30 and more at% of Mg do not react either with air or with water. In thermal decomposition of the hydrogenation product, H2 is sep-It is noted that $^{M}\mathcal{E}_{m}^{H}$ Ce H J r. These mixtures can be expressed arated in several stages by the formula MgCeH5. Both the method of hydrogenation of Ce-Mg alloys and the properties of the 'double hydrides" differ from the hydrogenation of Mg and Ce, and from the properties of CeH, and MgH, respectively. As to their properties, the products of complete hydrogenation of Je-Mg alloys (e. g., ard 2/4

30029 \$/020/61/141/001/013/021 B103/B147

Hydrogenation of cerium-magnesium...

MgH₂CeH₃ or MgCeH₅) take an intermediate position as compared to their constituents CeH₃ and MgH₂. There are 1 figure, 1 table, and 6 references: 4 Soviet and 2 non-Soviet. The reference to the English-language publication reads as follows: F. H. Ellinger, C. E. Holley et al., J. Am. Chem. Soc., 77, 2647 (1955).

AUGOCIATION: Institut obshchey i neorganicheskoy khimii im. N. S. Kurnakova Akademii nauk SSSR (Institute of General and Inorganic Chemistry imeni N. S. Kurnakov of the Academy of Sciences USSR)

TRE ENTED: April 29, 1961, by I. I. Chernyayev, Academician

SUBMITTED: April 24, 1961

Table 1. Composition of hydrogenation products of Ce-Mg alloys. Legend: (1) Composition of alloys; (2) hydrogen content; (3) number of test; (4) Mg, at%; (5) Mg, % by weight; (6) induction period, min; (7) time of hydrogenation; (8) per g of hydride, milliliters; (9) per g of Ce, Card 3/4

X

"APPROVED FOR RELEASE: 08/26/2000 CIA-RDP8

CIA-RDP86-00513R001653310007-5

MIKHEYEVA, V.I.; STERLYADKINA, Z.K.; KONSTANTINOVA, A.I.;

KRYUKOVA, O.N.

Absorption of hydrogen by alloys of cerium with magnesium.

Zhur. neorg. khim. 8 no.6:1314-1319 Je '63. (MIRA 16:6)

1. Institut obshchey i neorganicheskoy khimii imeni N.S.

Kurnakova AN SSSR, laboratoriya khimii gidridov i bora.

(Cerium-magnesium alloys)

(Hydrogenation)

MIKHEYEVA, V.I.; STERLYADKINA, Z.K.; CHERTKOV, A.A.

Hydrogenation of cerium alloys with aluminum. Zhur. neorg.
(MIRA 16:7)

khim. 8 no.7:1710-1714 Jl '63,

1. Institut obshchey i neorganicheskoy khimii imeni N.S.
Kurnakova AN SSSR, laboratoriya khimii gidridov i bora.
(Cerium-aluminum alloys) (Hydrogenation)

L 34499-65 EMP(e)/EWP(t)/EMP(b) IJP(c) JD ACCESSION NR: AP5002795 S/0078/65/010/001/0010/0017

AUTHOR: Sterlyadkina, Z. K.; Kryukova, O. N.; Mikheyeva, V. I.

194

TITLE: Reaction of potassium borohydride with sulfur

SOURCE: Zhurnal neorganicheskoy khimii, v. 10, no. 1, 1965, 10-17

TOPIC TAGS: potassium borosulfide, synthesis, potassium borohydride sulfur reaction, potassium borohydride

ABSTRACT: A thermodynamic study was made of the irreversible reaction of KBH₄ with S occurring upon heating mixtures of 0-100% of each to 700C. This and hydrogen evolution curves indicated 2 principal reactions: formation of KBS₂ at 230-300C: KBH₄ + 2S \longrightarrow KBS₂ + 2H₂(1), and thermal decomposition of the borohydride, KBH₄ \longrightarrow K + B + 2H₂(2). The most pure KBS₂ was obtained in highest yield when the S:KBH₄ molar ratio was 2. The deviation from theoretical in the yield of H₂ when either component was used in excess of this ratio indicated reactions other than (I): The series of thermal effects noted in the tempera-

Card 1/2

L 34499-65

ACCESSION NR: AP5002795

ture range between the beginning of reaction (1) and the beginning of the thermal decomposition of KBH4 was attributed to phase transformations involving participation of the reaction product KBS₂ and excess reactants -- KBH4 when S is in the 0--66.7 at.\$ range, and S when present in amounts above 66.7 at.\$. Orig. art. has:

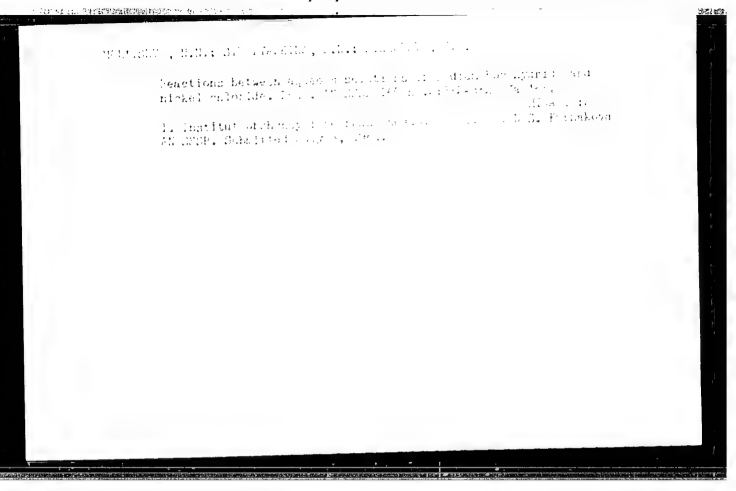
7 figures, 3 equations, and 2 tables.
ASSOCIATION: Institut obshchey i neorganicheskoy khimii im. N. S. Kurnakova
Akademii nauk SSSR (Institute of General and Inorganic Chemistry, Academy of

Sciences SSSR)

SUBMITTED: 01Jul63 ENCL: 00 SUB CODE: GC

NR REF SOV: 004 OTHER: 002

Card 2/2



"APPROVED FOR RELEASE: 08/26/2000

CIA-RDP86-00513R001653310007-5

EPF(c)/EPA(s)-2/ENT(m)/EWP(b)/EWP(e)/EWP(t) # Pr-4/Pt-7 L 43751-65 JD/JG ACCESSION NR: AP5008478 \$/0078/65/010/003/0583/0587 AUTHOR: Sterlyadkina, Z. K.; Kryukova, O. N.; Mikheyeva, V. I. CAMPACTURE CALFORNIA CONTRACTOR C B TITLE: Reaction of sodium borohydride with sulfur RUE: Zhurnal neorganicheskoy khimii, v. 10, no. 3, 1965, 583-587 TOPIC TAGS: sodium borohydride, potassium borohydride, sulfur, hydrogen sulfide, hydrogen, sodium borosulfide, sodium borosulfide production, gasometric analysis, - ...rmographic analysis, chemical analysis ABSTRACT: The reaction of sulfur with sodium borohydride has been studied to determine the production conditions for sodium borosulfide and hydrogen The experiments are carried out with NaBH4-S mixtures heated to 7500 and the results are compared with those obtained for reactions of KBH4-S mixtures. The thermographic, gasometric, and chemical analyses of the NaBH4-S mixtures show two basic reaction trends, viz., formation of sodium borosulride and hydrogen, and thermal decomposition of nonreactive sodium borohydride proceeding at elevated temperatures in mixtures with higher NaBH4 content. The formation of sodium borosulfide takes place at a lower temperature (225-230C) than the formation of potassium borohydride Card 1/42

"APPROVED FOR RELEASE: 08/26/2000

CIA-RDP86-00513R001653310007-5

L 43751-65

ACCESSION NR: AP5008478

(230-2500) and is accompanied by more side reactions, particularly the formation of hydrogen sulfide. The experimental results are given in Tables 1 and 2 of the Enclosure. The data obtained make it possible to assume that solid solutions of sodium borosulfide with excess hydrogen sulfide are formed from sodium borohydride and sulfur mixtures. Orig. art. has: 3 formulas, 4 figures, and 2 tables.

ASSOCIATION: none

SUBMITTED: 22Sep63

ENCL: 02

SUB CODE: GC

NO REF SOV: 006

Card

OTHER: 001

STERLYACKINA, C.K., KRYUKOVA, O.N., MIKHEYEVA, V.I.

Reaction of potentian borohydride with sulfur. Zhur. neorg. khim. 10 no.1:10-17 sa '65. (MIRA 18:11)

1. Institut obshchey i neorganicheskoy khimii imeni Kurnakova AN SSSR. Submitted July 1, 1963.

STERLYAUKHEA, Z.K.; KRYUKOVA, C.N.: MIEHEYEVA, V.1.

Resolvion of sedium boron hydride with sulfur. Zhur. neorg.
(MIEA 18:7)

khim. 10 no.3:583-587 Mr '65.

GUTKOVSKIY, V.A., kand.tekhn.nauk (Orsha); STERLYAGOV, A.A.;
MIKLASHEVSKIY, S.N., insh. (Orsha)

Highly efficient utilisation of steam locomotives. Zhel. dor.
transp. 40 no.3:70-72 Mr '58. (MIRA 11:4)

1. Hachal'nik depo Orsha, Belorusskoy dorogi (for Sterlyagov).
(Locomotives)

Activities of technology clubs and workshops in schools. Fig. v shkole 14 no.3:62-65 My-Je '54. (MLRA 7:7)

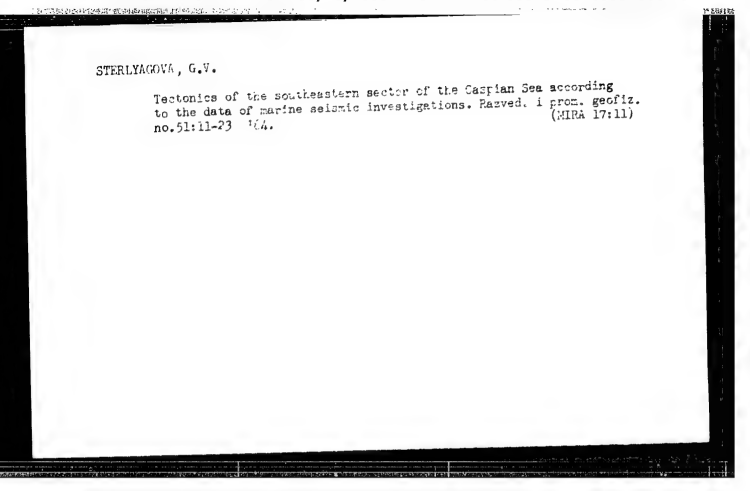
1. 569-ya sredwyn sakola, g. Moskva.
(Physics—Study and teaching) (Technology—Study and teaching)

CIA-RDP86-00513R001653310007-5

Papular scientific literature on the properties of solids.

Fiz. v shkole 15 no.3:85-86 My-Je '55.

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[Fishes and commercial fishing in Lake Baikal] Ryby i rybnoe khoziaistvo v basseine ozera Baikal. Irkutskoe, knizhnoe isd-vo, 1958. 745 p. (MIRA 12:4)

1. Sotrudniki Irkutskogo gosuniversitets (for Misharin, Tomilov, Popov, Yegorov, Tugarina). 2. Sotrudnik Baykal'skoy limnologicheskoy stantsii Akademii nauk SSSR (for Koryakov). 3. Baykalrybtrest (for Spelit, Artyunin). 4. Gosplan Buryat-Mongol'skoy ASSR (for Shniper). (Baikal, Lake--Fisheries)

